FORM PTO-1390 (REV 11-98) U S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE TRANSMITTAL LETTER TO THE UNITED STATE DESIGNATED/ELECTED OFFICE (DO/EO/US CONCERNING A FILING UNDER 35 U.S.C. 371 INTERNATIONAL APPLICATION NO PCT/GB99/00737 INTERNATIONAL FILING DATI 22 March 1999

EXPRESS MAIL EK802769739US

ATTORNEY'S DOCKET NUMBER

57.0272PCT

U.S. APPLICATION NO. (If known see 32 CFP 1.5 09/646/15

PRIORITY DATE CLAIMED 20 March 1998

TITLE O	F INVENTION HYDROPHOBIALLY MODIFIED POLYMERS FOR WATER CONTROL
APPLICA	NT(S) FOR DO/EO/US Timothy Gareth John JONES; Gary John TUSTIN
1. X 2. 3. X 4. X 5. X	herewith submits to the United States Designated/Elected Office (DO/FO/US) the following items and other information.  This is a FIRST submission of items concerning a filing under 35 U.S.C. 371.  This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371  This express request to begin national examination procedures (35 U.S.C. 371(f) at any time rather than delay examination until the expiration of the applicable time limit set in 35 U.S.C.371(b) and PCT Articles 22 and 39(1)  A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date.  A copy of the International Application as filed (35 U.S.C. 371(c)(2))  a.  is transmitted herewith (required only if not transmitted by the International Bureau).  b.  is not required, as the application was filed in the United States Receiving Ofice (RO/US).  A translation of the International Application into English (35 U.S.C. 371(c)(2)).  Amendments to the claims of the International Application under PCT Article 19(35 U.S.C. 371(c)(3))
8.	a.
	1. to 16. below concern document(s) or information included:
	An Information Disclosure Statement under 37 CFR 1.97 and 1.98.  An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.
13.	A FIRST preliminary amendment.
	A SECOND or SUBSEQUENT preliminary amendment.
14.	A substitute specification.
15.	A change of power of attorney and/or address letter.
16. X	Other items or information:

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Independent claims	_13 =	-0-	X \$78.00	\$ -	
MULTIPLE DEP	ENDENT CLAIM(S) (if appli		+ \$260.00	\$	
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		SUBT	TOTAL =	\$	
Processing fee of months from the	\$130.00 for furnishing the earliest claimed priority date	English translation later than te (37 CFR 1.492(f)).	20 30 +	\$	
		TOTAL NATION	AL FEE =	s 840.00	
Fee for recording accompanied by	the enclosed assignment (3 an appropriate cover sheet (	37 CFR 1.21(h)). The assignment of CFR 3.28, 3.31). \$40.00 pe	ent must be r property +	\$	
		TOTAL FEES ENC	LOSED =	\$ 840.00	
				Amount to be:	\$
				charged	\$
a. A check in the amount of \$ to cover the above fees is enclosed.					
b. A duplicate copy of this sheet is enclosed.  04-1579 in the amount of \$\\$840.00\$ to cover the above fees.					
c. The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 04-1579 A duplicate copy of this sheet is enclosed.					
NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the application to pending status.					
SEND ALL CORRESPONDENCE TO:				KITT	
Gordon G.			SIGNAT	URE:	
	ger Technology Corpora	on G. Waggett	~		
	berger Drive, MD1		NAME		
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Pn: 281-28	5-4524; Fax: 281-285-8	3569			
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# Hydrophobically Modified Polymers for Water Control

The present invention relates to cross-linkable hydrophobically modified water-soluble polymers for reducing the water

5 production of a hydrocarbon well drilled through subterranean formations. More particularly it relates to such polymers and polymer containing compositions for selectively reducing the flow of subterranean aqueous fluids into a well while maintaining the hydrocarbon production.

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### BACKGROUND OF THE INVENTION

During the life cycle of a hydrocarbon well, e.g., a well for
extracting oil or natural gas from the Earth, the producing well
commonly also yields water. In these instances, the amount of
water produced from the well tends to increase over time with a
concomitant reduction of hydrocarbon production. Frequently, the
production of water becomes so profuse that remedial measures
have to be taken to decrease the water/hydrocarbon production
ratio. As a final consequence of the increasing water
production, the well has to be abandoned.

Various techniques have been developed and used for reducing the quantity of water produced from oil and/or gas wells. For example, US Pat. No. 3,719,228 discloses a method of treating a subterranean formation containing hydrocarbons and brine to stimulate the production of hydrocarbons: A preflush composition comprised of a water solution of rosin soap and fatty acid soap is injected into the formation. The preflush reacts with connate brine to produce a precipitate that blocks the brine-bearing passages. The composition does not react with hydrocarbons thereby allowing hydrocarbon-bearing passages to remain open. Another alternative method is disclosed in US Pat. No.

4,617,132: A sandstone formation is contacted with an aqueous solution containing a water soluble anionic polymer having a

molecular weight greater than 100,000. As the next step, the

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anionic polymer is contacted with a fluid containing a water soluble cationic polymer having a weight greater than 1,000. As a result of the contact of the anionic with the cationic polymer, coacervation occurs between the two polymers which reduces the amount of the anionic polymer removed from the formation by fluids produced therefrom. The presence of stabilized polymer in the formation reduces the water/oil ratio by reducing the permeability of the formation to water in the wellbore area.

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In yet another technique, disclosed in US Pat. No. 5,203,834, comprises the steps of injecting a gas, a polymer composition capable to form a foamed gel with the gas, and a delayed gel degrading agent being capable of opening pathways in the gel.

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The gelation of high molecular weight polymers  $(M_w > 10^6 \text{ g/mol})$ has been extensively used in the development of water-based treatment fluids for water control is further described for example by R. D. Sydansk in "Acrylamide-polymer/chromium(III) -20 carboxylate gels for near wellbore matrix treatments", 7th SPE Symp. Enhanced Oil Recovery, Tulsa, Ok., April 1988, SPE/DoE 20214, or by R. S. Seright in: "Placement of gels to modify injection profiles", SPE/DoE Symp. Enhanced Oil Recovery, Tulsa, Ok., April 1994, SPE 27740. Typically for those methods, an 25 aqueous solution of a high molecular weight polymer, such as a polyacrylamide/polyacrylate copolymer (a so-called partiallyhydrolysed polyacrylamide), is gelled in situ in a porous formation using a metal cross-linker such as Cr3+ or small watersoluble organic cross-linkers such as formaldehyde and 30 formaldehyde/phenol. Other water-soluble polymers such as poly(vinyl alcohol), the polysaccharide guar gum and the copolymer poly(vinylpyrrolidone-co-2-acrylamido-2-methyl-1propanesulphonic acid) which can be cross-linked with a variety of cross-linking agents such as Zr4+ and boric acid.

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A more recent approach is described by A. Keller and K. A. Narh in: "The effect of counterions on the chain conformation of polyelectrolytes, as assessed by extensibility in elongational

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flow: the influence of multiple valency", J. Polym. Sci.: Part B: Polymer Phys., 32, 1697-1706 (1994). It includes the crosslinking of poly(sodium 4-styrenesulphonate) using Al3+ ions to form a gel. The concentration of the high molecular weight 5 hydrophilic polymers used to form hydrogels is typically in the range 3-10 g/l.

Hydrophobically-modified polymers as such and their application as thickening agents in aqueous fluids has been extensively 10 described. The principal application of these polymers is as viscosifying agents and they are commonly referred to as associative thickeners; these polymers are also known as polysoaps. The associative behaviour is generated by a small mole fraction (typically  $\leq 0.05$ ) of hydrophobic groups which, as separate monomers, are not soluble in water. The hydrophobic groups undergo intermolecular association in aqueous solution above the overlap concentration which results in a significant increase in solution viscosity.

20 Hydrophobically-modified polymers can be gelled in the presence of monomeric surfactants. For example, large increases in the viscosity of solutions of hydrophobically-modified poly(acrylic acid) in the presence of anionic, cationic and non-ionic surfactants have been disclosed. A brief review of the interaction between hydrophobically-modified polymers and surfactants can be found in: Lindman, B. and Thalberg, K., "Polymer-surfactant interactions - recent developments" in Interactions of Surfactants with Polymers and Proteins, E.D. Goddard and K.P. Ananthapadmanabhan (eds.), pp. 203-276, CRC Press, Boca Raton, Florida (1992).

In the U.S. Patent No. 4,532,052, the use of branched organic polymers for various well treating operations is described. The known polymers include hydrophilic and combinations of 35 hydrophilic and hydrophobic side groups. The preferred examples are characterized as having large hydrophilic branches.

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In the U.S. Patent No. 5,003,006, there are described crosslinked block copolymers containing polar and non-polar segments, with the polar segments generally making up at least 50 per cent.

It is the object of the invention to provide new polymers for water control application.

#### SUMMARY OF THE INVENTION

According to a first aspect of the invention, hydrophobically modified water-soluble polymers are provided which are capable of being chemically cross-linked to produce permanent gels in water-bearing formations. The polymers are characterized by a linear chain or backbone and attached thereto hydrophobic side groups at random locations. The term "linear" is defined as the absence of extended side groups/chains others than the hydrophobic groups. The term "extended" can be interpreted as having a molecular weight of more than 200.

It has been found that hydrophobically modified polymers when based on block copolymers, tend to form an inferior gel presumably due to presence of inhomogeneous gelation. In addition there is economic advantage when producing polymers with high molecular weight through a random co-polymerization process. The use of large polymers is advantageous as it reduces the polymer concentration required to achieve a desired level of viscosity. Within the scope of the present invention, a high molecular weight can be defined as a molecular weight in excess of 50,000, more preferably in excess of 100,000. A preferable upper limit for the size of the polymers is given by a molecular weight of 15 million, preferably 5 million, more preferably 500,000 or less.

The polymers preferably comprise low concentrations (0.5-5 mole percent) of hydrophobic monomers which randomly co-polymerise with the hydrophilic monomers to form random copolymers. The

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hydrophilic units of the polymer, which constitute preferably 95-99.5 mole percent of the polymer chain, contain preferably at least some small fraction of functional groups which can be chemically cross-linked to produce a gel. Alternatively, the hydrophobic groups themselves can contain chemical functions which can be the sites of cross-linking. Preferred hydrophobic groups are alkyl groups of more than six , preferably eight, and even more preferably 12 carbon atoms, and derivatives thereof.

10 Chemical cross-linking is defined as forming a chemical bond between the cross-linked polymers. Chemical cross-linking is understood to be stable and thermally irreversible. Some of the advantages in the use of hydrophobically modified polymers are responsiveness to hydrocarbons resulting in a reduction in the adhesive strength of gels and an extreme retardation of gelation, and solubilisation of large organic cross-linking agents.

The cross-linking agents can be either inorganic ions (or ionic 20 complexes) or polar organic molecules. When the polymer contains ionic groups such as carboxylate or sulphonate functions the polymer chains can be cross-linked by inorganic ions such as chromium(III) or zirconium(IV), frequently in the presence of ligands, such as acetate or adipate ions, to control the rate of 25 cross-linking. Alternatively, organic cross-linking agents can be used. For example, the amide groups on poly(acrylamide) can be cross-linked with formaldehyde or a mixture of formaldehyde and phenol. The disadvantages of using formaldehyde or similar small molecules are their toxicity and issues of disposal. The 30 larger reagents such as hexanal and heptanal, which are not classed as toxic, would be suitable cross-linking agents but are not sufficiently soluble in water to cross-link normal hydrophilic polymers such as the poly(acrylamide)/poly(acrylate) copolymer. However, these larger cross-linking agents are 35 solubilised in aqueous solutions by hydrophobically-modified poly(acrylamide) polymers and can thence cross-link them.

These and other features of the invention, preferred embodiments and variants thereof, and further advantages of the invention will become appreciated and understood by those skilled in the art from the detailed description following below.

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### MODE(S) FOR CARRYING OUT THE INVENTION

A large number of hydrophobically-modified polymers are known as such and suitable for the purpose of the present invention. These polymers are based on the modification of common water-soluble (hydrophilic) polymers such as poly(acrylamide), poly(acrylic acid), poly(vinylpyridine), hydroxyethylcellulose and poly(ethylene oxide). The structure of examples of hydrophobically-modified polymers are given below for poly(sodium 4-styrenesulphonate (a), poly(vinylpyridine)(b), and poly(acrylamide)(c):

$$(a) \begin{array}{c} H \\ CH_2 - C \\ CH_2 - C \\ CH_3 \\ Na \end{array} \begin{array}{c} H \\ CH_2 - C \\ CH_3 \\ CH_4 \\ CH_3 \\ CH_4 \\ CH_5 \\ CH_$$

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(b) 
$$CH_2$$
  $CH_2$   $CH_$ 

$$\begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \\ \\ \end{array} \end{array} \end{array} \begin{array}{c} \begin{array}{c} \\ \\ \end{array} \end{array} \begin{array}{c} \\ \\ \end{array} \begin{array}{c} \\ \\ \end{array} \end{array} \begin{array}{c} \\ \\ \end{array} \begin{array}{c} \\ \\ \end{array} \begin{array}{c} \\ \end{array} \begin{array}{c} \\ \\ \end{array} \end{array} \begin{array}{c} \\ \\ \end{array} \begin{array}{c} \\ \\ \end{array} \begin{array}{c} \\ \end{array} \begin{array}{c} \\ \end{array} \begin{array}{c} \\ \\ \end{array} \begin{array}{c} \\ \end{array} \end{array} \begin{array}{c} \\ \end{array} \begin{array}{c$$

The fraction M/(M+N) denotes the mole fraction of hydrophobic monomers in the polymer.

The following formulae describe the structure of further examples which can be readily cross-linked using known crosslinkers.

The structure of a hydrophobically-modified poly(acrylamide) where the hydrophobic modification takes the form of replacing about 3 mole percent (M/(M+N)=0.03) of the acrylamide units with the hydrophobic monomer n-nonyl acrylate is:

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This polymer has an average molecular weight of about  $5 \times 10^6$ g/mole and it can be readily cross-linked with chromium(III) or zirconium(IV) or the organic cross-linker hexanal. Where the 20 gels formed by conventional poly(acrylamide)/poly(acrylate) copolymers undergo syneresis and shrink, the gels formed by hydrophobically-modified polymers in otherwise identical formulations undergo syneresis at approximately half the rate of the hydrophilic polymers. It is also observed that the crosslinking of the hydrophobically-modified poly(acrylamide) is less sensitive to the concentration of divalent ions  $(Ca^{2+}, Mg^{2+})$ .

The structure of another hydrophobically-modified poly(acrylamide) where the hydrophobic modification takes the form of replacing about 1 mole percent (M/(M+N)=0.01) of the acrylamide units with the hydrophobic monomer N-decylacrylamide

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is:

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$$(e) \qquad \begin{array}{c|c} & & & \\ \hline & & \\ & & \\ \hline & & \\ & & \\ \hline \end{array} \qquad \begin{array}{c} & & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ & \\ \end{array} \qquad \begin{array}{c} & \\ & \\ \end{array} \qquad \begin{array}{c}$$

This polymer can also be cross-linked using metal ions (Cr(III), Zr(IV)) and organic molecules (hexanal).

An additional feature of hydrophobically-modified polymers such as (d) and (e) is that they can be prevented from cross-linking in contact with hydrocarbon. The prevention of cross-linking can arise from two possible mechanisms. Firstly, the larger organic cross-linker can be removed from the polymer solution by solubilisation in the contacting hydrocarbon. The cross-linker hexanal, which is solubilized by the hydrophobically-modified polyacrylamide, is readily soluble in hydrocarbon and can be extracted from the polymer solution. Secondly, the solution of hydrophobically-modified polymer can partly solubilize the hydrocarbon which itself can be sufficient to prevent cross-linking. Thus a solution of hydrophobically-modified poly(acrylamide) with the metal ion cross-linker such as Cr(III) or a small water-soluble cross-linker such as formaldehyde can be prevented from cross-linking when mixed with hydrocarbon.

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#### CLAIMS

A composition for selectively blocking water bearing subterranean formations, said composition comprising water
 soluble polymers having a linear hydrophilic backbone with hydrophobic side groups located at random positions along said backbone and functional groups for cross-linking said polymers, said composition forming a cross-linked gel in a subterranean water-bearing formation.

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- 2. The composition of claim 1, wherein the polymers comprise 0.5 to 5 mole per cent of hydrophobic side groups.
- 3. The composition of claim 1, wherein the functional groups for cross-linking form part of the hydrophilic backbone of the polymer.
  - 4. The composition of claim 1, wherein the functional groups for cross-linking form part of the hydrophobic side groups.
  - 5. The composition of claim 1, wherein the polymers have a molecular weight of 50,000 or more.
- 6. The composition of claim 1, further comprising a chemical cross-linking agent.
  - 7. The composition of claim 6, wherein the chemical crosslinking agent is organic.
- 30 8. The composition of claim 6, wherein the chemical cross-linking agent is an aldehyde or aldehyde derivative comprising at least 5 carbon atoms.
- 9. The composition of claim 8, wherein the chemical crosslinking agent is hexanal or heptanal.

Patent and Trademark Office: U.S. DEPARTMENT OF COMMERCE

## **DECLARATION FOR PATENT APPLICATION**

Declaration Submitted	d <b>WITH</b> Initial	Filing	OR [	☑ Declarati	on Submitted	After Initial Filing
Attorney Docket Numb First-Named Inventor Application Number: Filing Date: Group Art Unit: Examiner's Name:		COMPLETE	IF KNOW	IN:	57.0272 Gareth Joh 09/646,715 ember 20, 2	
As a below-named	inventor, I hereby	declare that:				
My residence, post	office address and	citizenship a	re as stated	below next to	my name.	
inventor (if plural names are invention entitled:		e subject ma	tter which	is claimed and	for which a pa	
the specification of which:						
is attached hereto as Atto	rney Docket No.:					
OR						
■ was filed onSeptember     □ PCT International Application     □ and was amended on	cation No			tes Application	n No. 09/646,71	5 or
I hereby state that the claims, as amended by any			the conten	its of the above	e identified spec	rification, including
I acknowledge the	duty to disclose in	formation wh	ich is mate	rial to patental	oility as defined	in 37 CFR § 1.56.
I hereby claim application(s) for patent or i least one country other than foreign application for patent the application on which prior	nventor's certificate the United States, or inventor's certif	e, or § 365(a listed below	of any Po and have	CT Internation also identified	al application v d below, by che	ecking the box, any
Prior Foreign Application Numbers	Country		iling Date DYY)	Priority N Claimed		Copy Attached? No
9805880.3	GB	•	AR-98			×
PCT/GB99/00737	WO		AR-99			
Additional foreign applie	cation numbers are	listed in a su	pplemental	priority data s	heet PTO/SB/02	2B, attached hereto.

☐ I hereby claim the benefit under Title 35 U.S.C. § 1.19(e) of any United States provisional application(s) listed below.

Application Number	Filing Date

Additional provisional patent application numbers are listed in a supplemental priority data sheet PTO/SB/02B, attached hereto.

I hereby claim the benefit under 35 U.S.C. § 1.20 of any United States application(s), or § 365(c) of any PCT International application designating the United States, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph 35 of U.S.C. § 112, I acknowledge the duty to disclose information which is material to patentability as defined in 37 CFR § 1.56 which became available between the filing date of the prior application and the national or PCT International filing date of this application.

<b>US Parent</b>	PCT Parent Number	<b>Parent Filing Date</b>	Parent Patent Number
Application Number		(MMDDYY)	(if applicable)

Additional US or PCT international application numbers are listed in a supplemental priority data sheet PTO/SB/02B, attached hereto.

As a named inventor, I hereby appoint the following attorney(s) and/or agents(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith:

Name	Registration Number	Name	Registration Number
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Douglas Y'Barbo	42,239	William L. Wang	39,871
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Additional registered practitioner(s) named on supplemental Registered Practitioner Information Sheet PTO/SB/02B, attached hereto.

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I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the

United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Sole or first inventor:			
Inventor's Full Name	Timothy	<u>,G.J.</u>	<u>Jones</u>
	(First)	(Initial)	(Last)
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Country of Residence:	U.K.	Citizenship:	U.K.